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ETHYLENE (CO)POLYMERS, AND LAMINATE AND HOLLOW MOLDING MADE BY USING THE SAME:

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ABSTRACT:

Ethylene (co)polymers each having specified values of (1) density, (2) molecular weight distribution (Mw/Mn), (3) relationship between die swell (DS) and shear rate (gamma) in a region of shear rate (gamma), (4) relationship between stress (sigma) and maximum stress (sigma max) in a stress-strain curve in uniaxial stretching, (5) the ratio of sample diameter to initial sample diameter and the time for sample breakage during uniaxial stretching under a constant tension, laminates and hollow moldings made by using the same. The ethylene (co)polymers are excellent in drawdown resistance, puncture resistance, parison control response, pinch-off properties, durability, and fire resistance.

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(54) ETHYLENE (CO)POLYMERS, AND LAMINATE AND HOLLOW MOLDING MADE BY USING THE SAME

(57) Ethylene (co)polymers each having specified values of (1) density, (2) molecular weight distribution (MwMm), (3) relationship between die swell (DS) and shear rate (r) in a region of shear rate (r), (4) relationship between stress (r) and maximum stress (crimation as stress-strain curve in uniaxial stelching, (5) the ratio of sample diameter to initial sample diameter and the time for sample breakage during uniaxial stelching under a constant tension, laminates and hollow moldings made by using the same. The ethylene (oppolymers are excellent in drawdown resistance, puncture resistance, parison control response, pinch-off properties, durability, and fire resistance.





(a)

Description

TECHNICAL FIELD

This invention relates to a novel ethylene (co)polymer, and laminate and hollow molding or blow molded products made by using the same. More particularly, this invention relates to an ethylene (co)polymer which is excellent in mechanical strengths such as impact resistance, moldability, etc. and applicable to extrusion molding, injection molding, blow molding, etc., which is excellent for use in large blow molded products required of drawdown resistance, puncture resistance, parison thickness controllability (parison control response), pinch-off properties, durability, etc., in particular excellent in fire resistance required for fuel tanks, to hollow moldings, laminates and hollow multilayer moldings made by using the same.

BACKGROUND ART

- Generally, detergent bottles, food containers such as beverage bottles and vegetable oil bottles, large vessels such as drum cans and industrial cans, fuel containers such as kerosene cans and gasoline tanks, spoilers, bumpers and the like are produced by a blow molding process. The blow molding process is to pinch with a mold a cylindrically extruded perison composed of a melted resin, blowing air into the perison to blow up and change the shape of it so as to conform to the shape of the cavity of the mold, and then cooling it.
- The blow molding process is widely utilized since it is applicable widely to large fuel tanks having complicated shapes, drums, and further panel-like moldings as well as hollow moldings such as bottles and since molding is simple and molding costs covering molds are inexpensive.
 - Recently, in the field of large vessels such as drum cans and large containers, substitution by plastics is being carried forward positively taking into consideration freedom in shape, economy, safety, and adaptability to envi-
- It is also the case in the field of automobile industry such as fuel tanks, bumpers and spoilers. Particularly in 25 ronmental problems. the field of fuel tanks, substitution by plastics proceeds remarkably along with the development of fuel permeation preventing technology by a co-extrusion technology, a sealer technology, a continuous multi-ply technology, etc.
- When large products are blow molded, a phenomenon that a parison droops down due to its own weight 30 (drawdown) tends to occur. To make the drawdown small, it has been known to use a resin having sufficiently high vis-
 - In molding hollow moldings having a complicated shape, there tends to occur a local increase in blow ratio cosity and melt tension. to cause puncture or local thinning of a parison. To prevent the puncture or local thinning of a parison, it has been known to control the thickness of the parison (parison control) to make a desired portion thick or make alterations in the appli-
- 35 ance such as alteration of the shape of a mold. Conventionally, there has been a problem that an increase in molecular weight and an increase in viscosity in order to improve the drawdown resistance of polyethylene results in aggravation of extrusion properties (extrusion rate, parison surface conditions) and welded strength of a parison becomes worse so that the pinch-off shape is worsened. As a method for solving the problem have been known, for example, the method in which a multi-step polymeri-2 zation method using a Ziegler-type catalyst (Japanese Patent Application Laid-open No. Sho 55-152735), the method
 - in which a small amount of radical generator and crosslinking auxiliary are added to polyethylene resin (Japanese Patent Publication Hei 2-52654), the method in which two polyethylene components are blended in a certain proportion (Japanese Patent Application Laid-open No. Hei 6-299009), etc. However, recently larger and more complicated blow moldings are being produced so that further improve-
- 45 ment of drawdown resistance of a resin is becoming necessary in order to extrude a heavy parison to a larger length in a stable manner. Also, in order to improve the fuel permeation of a fuel tank, there is a demand for a multilayer fuel tank provided with an ethylene/vinyl acetate copolymer seponification product (hereafter, called EVOH), which has poor heat resistance, as a barrier material. In molding them, a blow molding machine of a continuous extrusion type having a smaller retention portion and suffering less thermal decomposition of EVOH is becoming to be put in use for blow mold
 - ing larger fuel tanks in place of a blow molding machine of an accumulator type which has heretofore been used for larger size molding. As compared with the accumulator type, the continuous extrusion type molding machine takes a long time for extruding a parison so that drawdown of a parison tends to occur more frequently. Therefore, there is a demand for a resin having an increased drawdown resistance.
- [0010] For this purpose, the improved polyethylenes obtained, for example, by the above-mentioned processes are 55 insufficient in drawdown resistance for obtaining large hollow moldings.
 - [0011] Also, according as the shape of hollow moldings becomes more complicated, there come into question phenomena that parisons will puncture when they are being formed by blowing and that local thinning of a product will occur at its corner portions, etc.

To prevent these problems, a method has generally been used in which a parison controller is used in order to control the thickness of a parison. However, conventional polyethylene and resin compositions thereof grow to have insufficient parison thickness controllability (parison control response) so that improvement of parison control response is desired. Also, as the method for enabling molding of deep drawn articles having complicated shapes, there have been known, for example, the method in which a split mold is used (Plastics, Vol. 42, No. 5, p.64-71) and the method in which a blow molding machine with a mold slanting mechanism is used (Plastics, Vol. 41, No. 10, p.59-69). However, the above-mentioned methods have a problem that costs for the apparatus and mold are high.

Furthermore, in the field of fuel tanks, which require fire resistance as one of the requisite properties, there have been proposed ethylene copolymers having an α-clefin content within a specified range and inherent viscosity, zero shear viscosity and time for melt stretching breakage being prescribed to be within respective specified ranges as a preferred material (Japanese Patent Application Laid-open No. Hei 7-101433). However, they are in an insufficient state in molding processability and fire resistance to cope with weight reduction and thickness reduction.

DISCLOSURE OF THE INVENTION

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An object of the present invention is to overcome the defects of conventional polyethylenes and provide an ethylene (co)polymer having excellent drawdown resistance, puncture resistance, parison control response, and pinch-

Another object of the present invention is to provide blow molded products which are improved against the off properties. phenomena that the parison of blow molding having a complicated shape will puncture when expansion shaped and that there will occur local thinning of corner portions of products, by use of the above-mentioned ethylene (co)polymer

Still another object of the present invention is to provide blow molded multilayer containers having a capability of preventing permeation of various fuels and causing no local thinning.

[0017] In a first aspect, the present invention relates to an ethylene (co)polymer which satisfies the following requirements (a) to (d):

(a) a density of 0.93 to 0.98 g/cm3,

(b) a molecular weight distribution (Mw/Mn) of 25 to 50. (c) that values of slope a and of intercept to obtained from a die swell (DS) and a shear rate (r) measured in a region of shear rate (y) (6.08 to 24.8 sec 1) at 230 °C, in accordance with the equation (1) below 30

$$DS = a \times Ln(\gamma) + b$$
 (1)

(d) that the stress ratio of a stress (d) at a strain of 1.0 to a maximum stress (dmax) in a stress-strain curve for uniaxial stretching at 0.1 sec 1 measured at 170 °C is such that σmax/σ≥2.1.

Preferably, the present invention relates to an ethylene (co)polymer which further satisfies the requirement (e) that a value of (α) as defined by the equation (2) below

$$\alpha = \epsilon \hbar$$
 (2)

(wherein α is $\epsilon't$ when ϵ defined by ϵ = -Ln(sample diameter after t seconds 1,/initial sample diameter i_0) is 0.6) when uniaxially stretched under a constant tension such that initial stress (= tension/initial cross section of sample) measured at 170 °C is 3.0×10^4 Pa is not greater than 0.03 and time for sample breakage is not shorter than 25 sections.

More preferably, the present invention relates to an ethylene (co)polymer which further satisfies the requireonds.

(f) HLMFR \leq 10g/10min and which has a C₃₋₂₀ α -olefin content of 10 mol% or less.

In a second aspect, the present invention relates to blow molded products comprising the above-mentioned ethylene (co)polymer or composition thereof, preferably a fuel tank which exhibits superiority of the above-mentioned ethylene (co)polymer or composition thereof.

In a third aspect, the present invention relates to a laminate comprising a layer comprising the above-men-55 tioned ethylene (co)polymer or composition thereof, a barrier layer, and optionally an adhesive layer and preferably to a laminate in which the barrier layer comprises at least one member selected from polyamide resins, ethylene/vinyl acetate copolymer saponification products, polyester resins, poly/inylidene chloride resins and compositions thereof. It is desirable that for the adhesive layer, adhesive resins which comprise copolymers of unsaturated carboxylic acids or derivatives thereof with olefins or polyolefin resins modified with unsaturated carboxylic acids or derivatives thereof be

In a fourth aspect, the present invention relates to a blow molded multilayer container comprising a laminate which comprises a layer comprising the above-mentioned ethylene (co)polymer or compositions thereof, an adhesive layer and a barrier layer, and preferably such exhibits superiority in multilayer blow molded fuel tanks.

[0021] In a fifth aspect, the present invention relates to a blow molded multilayer container comprising a laminate which comprises at least one resin layer selected from the recycled resin composed of the above-mentioned laminate, or compositions of the ethylene (co)polymer according to the first aspect containing the recycled resin, or compositions of these with polyoletin resin, a barrier layer, and optionally an adhesive layer, and preferably such exhibits superiority 10 in physical properties and economy in blow molded multilayer fuel tanks.

BRIEF DESCRIPTION OF THE DRAWINGS

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Fig. 1 is diagrams illustrating pinch-off properties when blow molding ethylene (co)polymers (2) of example and comparative example using a mold (1), with (a) being a vertically cut cross sectional view with respect to the pinchoff line of blow molding, (b) and (c) being partial enlarged views of (a) at a pinch-off middle portion (M), with (b) showing that the pinch-off shape is in a good state and (c) showing that the pinch-off shape is in a bad state.

DETAILED DESCRIPTION OF THE INVENTION

Hereafter, the present invention will be described in more detail.

The present invention has been achieved based on the finding that ethylene (co)polymer comprising an ethzz ylene homopolymer or a copolymer of ethylene and α -olefin having 3 or more carbon atoms and having a density, a molecular weight distribution, a relationship between a die swell (DS) and a shear rate (γ), a stress ratio, a time for melt breakage of resin which are adjusted to specified ranges is excellent in drawdown resistance, puncture resistance, partson control response, pinch-off shape properties, fire resistance, etc.

The ethylene (co)polymer of the present invention is an ethylene homopolymer or a copolymer of ethylene 30 with an α-olefin having 3 to 20 carbon atoms. The α-olefin include, for example, propylene, 1-butene, 1-pentene, 1-hexene, 4-methylperatene-1, 1-octane, 1-decene, etc. The content of a-ofetin is 10 mal% or less. If the content of a-ofetin exceeds 10 mol%, there is the fear that the rigidity of (co)polymer decreases.

The ethylene (co)polymer of the present invention has a density of 0.93 to 0.98 g/cm³, and preferably 0.944 to 0.97 g/cm³. If the density is below 0.93 g/cm³, the rigidity and fire resistance decrease. On the other hand, if the den-

[0027] The ethylene (co)polymer of the present invention has a molecular weight distribution (MwMn) which must 35 sity exceeds 0.98 g/cm³, the impact strength decreases. be in the range of 25 to 50, preferably 27 to 50, and more preferably 30 to 50.

[0028] If the molecular weight distribution is below 25, drawdown resistance and extrusion properties are poor while a molecular weight distribution exceeding 50 results in an increased die swell (DS) and an increased slope (a) of the equation (1) described below, a decreased parison control response, and an increased smoke when extruding part-

The ethylene (co)polymer of the present invention requires that values of slope a and of intercept b obtained sons. from a die swell (DS) and a shear rate (y) measured in a region of shear rate (y) (6.08 to 24.8 sec 1) at 230 °C, in accordance with the equation (1) below

$$DS = a \times Ln(\gamma) + b$$
 (1)

(wherein Ln (γ) is a natural logarithmic value of shear rate (γ)) are such that a≤0.05 and b≤1.45. The value of slope a is preferably in the range of a=0.03, and the value of intercept b is preferably in the range of b=1.35.

50 (1030) Where the value of slope a is a \$0.05 and the value of intercept b is to 1.45, or the value of slope a is a>0.05, control of the thickness of a parison (parison control response) becomes difficult.

[0031] The ethylene (co)polymer of the present invention requires that the stress ratio of a stress (r) at a strein of 1.0 to a maximum stress (ormax) in a stress-strain curve in unlaxial stretching at 0.1 sec 1 measured at 170 °C is in the range of omax/o≈ 2.1, and preferably 2.5 or more. Where the stress ratio (omax/o) is less than 2.1, the puncture resist-

[0032] More preferably, the ethylene (co)polymer of the present invention requires that a value of (a) as defined by the equation (2) below

(2) .

(wherein α is εt when ε defined by $\varepsilon = -Ln(sample diameter after t seconds <math>1_{\ell}$ finitial sample diameter 1_0) is 0.6) when unlaxially stretched under a constant tension such that initial stress (= tension/initial cross section of sample) measured at 170 °C is $3.0x10^4$ Pa is not greater than 0.03 and time for sample breakage is not shorter than 25 seconds. If α exceeds 0.03, the deformation speed is higher whereas time for breakage less than 25 seconds results in a shortened hole opening time, so that there is the fear that fire resistance will be poor.

The ethylene (co)polymer of the present invention desirably has (f) HLMFR (High Load Melt Flow Rate) of 10.0 g/10 min or less. In particular, HLMFR for large blow molded products such as fuel tanks is preferably 5.0 (g/10 min) or less, and more preferably 4.0 (g/10 min) or less. Where HLMFR is above 10.0 (g/10 min), there is the fear that

the drawdown resistance will remain unimproved. [0034] The ethylene (co)polymer of the present invention is excellent in molding processability, rigidity and mechanical properties and is applied to extrusion molding, injection molding, blow molding, etc. Satisfaction of the above-mentioned requirements (a) to (e), and further the requirement (f) in the case of large or complicated-configured moldings such as tanks, overcomes the delects that the conventional polyethylene for blow molding has and provides the moldings with superiority in drawdown resistance, puncture resistance, parison control response, pinch-off properties or fire

The method for producing the ethylene (co)polymer of the present invention is not limited particularly so long resistance. as the requirements (a) to (d) above, desirably further the requirement (e), and more desirably further the requirement (f), are satisfied and it can be produced by a gas phase polymerization method, a slurry polymerization method, a solution polymerization method, etc., in the presence of a catalyst such as a Ziegler type catalyst, a Phillips type catalyst, a metallocene type catalyst or the like. Polymerization conditions are not limited particularly, but the polymerization temperature is usually 15 to 350 °C, preferably 20 to 200 °C, and more preferably 50 to 120 °C. The polymerization pressure is usually from atmospheric to 70 KgG/cm², preferably atmospheric to 50 KgG/cm² in the case of low/medium pressure 25 methods, whereas in the case of high pressure methods, it is desirable that the polymerization pressure is usually not higher than 1,500 KgG/cm². The polymerization method may be one step polymerization or multi-step polymerization having two-steps or more in which different polymerization conditions such as hydrogen concentration, monomer concentration, polymerization pressure, polymerization temperature, catalyst, etc. are used and is not limited particularly. It is also possible to blend a plurality of components having different properties to prepare the ethylene (co)polymer of

In the present invention, known additives such as antistatic agents, antioxidants, lubricants, anti-blocking 30 the present invention. agents, anti-fogging agents, organic or inorganic pigments, fillers, ultraviolet absorbents, dispersants, weathering agents, crosslinking agents, toaming agents, flame retardants, etc. can it desired be added in such amounts that the

The hollow moldings of the present invention are hollow molding made of the above-mentioned ethylene object is not obstructed. (co)polymer or composition thereof and include detergent bottles, food containers such as beverage bottles and vegetable oil bottles, vessels such as lerosene cans, drums and medicine bottles, various fuel containers, various components such as spoilers, bumpers, etc., housings of home electric products, OA apparatus, etc., panel-like construction components for constituting simple toilets, pallets and the like.

The above-mentioned ethylene (co)polymer or composition thereof may be blended with in addition to the ethylene (co)polymer of the present invention other polyclefin resin in amounts of 1 to 70% by weight. The method of producing hollow moldings of the present invention includes an extrusion blow molding method, an injection blow molding method, an extrusion stretching blow molding method, an injection stretching blow molding method, etc., but not limited particularly to a cold parison method, a hot parison method, etc.

In another aspect, the present invention relates to a laminate, comprising at least a layer comprising the above-mentioned ethylene (co)polymer or composition thereof (hereafter, referred to as the present resin layer), and a barrier layer. The above-mentioned laminate is not limited particularly so long as there is a structure of the above-mentioned two types of layers and the order of their lamination is not limited particularly. Generally, it is preferred that the barrier layer is an intermediate layer and along with (an) adhesive layer(s) an inner layer and/or outer layer is constituted by the present resin layer, and the outer or inner layer is an other polyolefin layer.

The barrier layer used in the present invention is constituted by at least one resin selected from polyamide resins, ethylene/vinyl acetate copolymer saponification products, polyacrylonitrile resins, polymethacrylonitrile resins, polyacetal resins, polyester resins, polyvinylidene chloride resins, polycarbonate resins, or compositions thereof. From the viewpoint of performance and physical properties, polyamide resins, ethylenevinyl acetate copolymer saponificass tion products, poly(meth)acrylonitrile resins, polyester resins and polyvinylidene chloride resins are desirable.

The polyamide resins include polyamides obtained by polycondensation of a diamine with a dicarboxylic acid, polyamides obtained by condensation with an aminocarboxylic acid, polyamides obtained from a lactam, or copolymerized polyamides therefrom and those having a relative viscosity in the range of 1 to 6, a melting point of 170 to 280 °C, preferably 200 to 240 °C are used. More specifically, there can be cited nylon-6, nylon-66, nylon-610, nylon-9, nylon-11, nylon-12, nylon-666, nylon-66/150, nylon-671 and mixtures or polymer alloys of these polyamide resins with polyoletin resins and the like. Of these, nylon-6 is preferred.

[0042] The above-mentioned ethylene/vinyl acetate copolymer saponification products include, for example, ethylene/vinyl acetate saponification products such as an ethylene/vinyl acetate copolymer having a degree of saponification of 90% or more, preferably 95% or more and an ethylene content of 15 to 50 more.

10043] The poly(meth)acrylonitrile resins are (co)polymers which are polymers composed of a methacrylonitrile [10043]. The polymers composed of these units and an alkyl (methacrylate unit, with the ratio by unit and/or acrylonitrile unit or copolymers composed of these units and an alkyl (methacrylate unit, with the ratio by unit and/or acrylonitrile unit or copolymers composed of these units and an alkyl (methacrylate unit, with the ratio by weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average molecular weight measured by a GPC weight being in the range of 95.5 to 70.30, and which have a weight average of 95.5 to 70.30, and which have a weight average of 95.5 to 70.30, and which have a weight average of 95.5 to 70.30, and which have a weight average of 95.5 to 70.30, and which have a weight average of 95.5 to 70.30, and which have a weight average of 95.5 to 70.30, and which have a weight average of

method in the range of 60,000 to 200,000.
[0044] The above-mentioned polyester resins include thermoplastic polyesters such as polyethylene terephthalate, polyethylene terephthalate, byolyethylene terephthalate, polyethylene naphthalate, and benzoic acid polyester and mixtures or polymer alloys of these with polyoletins.

[0065] The above-mentioned polyvinylidene chloride resins are polymers having a vinylidene chloride unit content of, for example, 70 to 99 mol%, and preferably 80 to 98 mol% and examples of componener of the polyvinylidene chloride resin include acrylic monomers such as acrylonitrile, methacrylonitrile, methyl methacrylate, ethyl methacrylate, ethyl methacrylate, glycidy methacrylate, glycidy acrylate, hydroxyethyl acrylate, and hydroxypropyl acrylate, and independent such as wind chloride.

vinyl monomers such as vinyl chloride.

[0046] The adhesive layer used in the present invention is constituted by a resin which is at least one adhesive resin [0046] The adhesive layer used in the present invention is constituted by a resin which is at least one adhesive resin selected from copolymers of an unsaturated carboxylic acid or derivatives thereof grafted thereto or resin composition which is a mixtures of that adhesive resin with unmodified polyoletin or the like.

[0047] Specific examples of the above-mentioned copolymers of unsaturated carboxylic acid or derivative thereof and an otelin include binary or more copolymers such as ethylene/(meth)acrylic acid copolymers, ethylene/glycidyl (meth)acrylate copolymers, ethylene/glycidyl (meth)acrylate/inql acetale terpolymers, ethylene/melaic anthydride copolymers, ethylene/ethylacrylic acid/melaic anthydride terpolymers, styrene/melaic anthydride terpolymers, and ethylene/styrene/melaic anthydride terpolymers.

[0048] The polyoletin modified with an unsaturated carboxylic acid or derivatives thereof of the present invention are polymers obtained by graft modifying polyoletin with an unsaturated carboxylic acid or derivatives thereof in the presence of a free radical generator such as an organic percode in an extruder or a kneader such as a Henschel mixer,

or in a solvent such as a hydrocarbon.

[0049] Specific examples of the modified polyoletin include maleic anhydride-modified high density polyethylene, maleic anhydride-modified finear low density polyethylene, maleic anhydride-modified very-low density polyethylene, maleic anh

maleic anhydride-modified polypropylene and the like.

10050] These are used singly or as modified polyplefin compositions by blending them with the above-mentioned to make the polyplefin and optionally rubbers, etc., if desired.

[0051] The blending proportion of the adhesive resian and unmodified polyoletin is not limited particularly but generally the unmodified polyoletin may be blended in a blending proportion of 1 to 99% by weight. Further, it is desirable that the content of unsaturated carboxylic acid or derivatives thereof in the adhesive resin or composition thereof be selected in the range of 10⁻⁹ to 10⁻⁹ mole, and preferably 10⁻⁹ to 10⁻⁴ mole, per g of the resin component. With a content of below 10⁻⁹ mole, the adhesive strength is insufficient but a content exceeding 10⁻⁹ mole would cause "burn marks" to below 10⁻⁹ mole, the adhesive strength is insufficient but a content exceeding 10⁻⁹ mole would cause "burn marks"

[0052] The above-mentioned polyoletin resins include high/medium density polyethylenes, linear low density polythylene, very-low density polyethylene, low density polyethylene by high pressure radical polymerization, ethylethylene, very-low density polyethylene, low density polyethylene by high pressure radical polymerization, ethylethylene/unsaturated carboxylic acid allyle steet copolymers such as ethylene/ethyl (meth)lacrylate copolymers, propyene homopolymers, propylene/a olefin random or block copolymers, colefin homopolymers such as t-butene polyenes, etc., mutual oppolymers of these or the like. Of these, high/medium density polyethylenes, linear low density
polyethylene, very-low density polyethylene, etc. are preferred.

[0053] The unsaturated carboxylic acid or derivatives thereof used in the present invention include acrylic acid, methacrylic acid, maleic acid, furnario acid, itaconic acid, citraconic acid or their anhydrides, metal salts and the like. Of these maleic anhydride is orderered.

[0054] The free radical generator used for graft modification includes organic peroxides, dicumyl compounds, etc. Because of reactivity and ease of handling, organic peroxide-base compounds are preferred. Specific examples the reof include dicumyl peroxide, 2,5-dimethyl-2,5-bis(t-butylperoxy)beynes,1,3-bis(24-butylperoxy)seproxy)beynes,1,3-bis(24-butylperoxy)seproxy)beynes,1,3-bis(24-butylperoxy)seproxy)beynes,1,3-bis(24-butylperoxy)seproxy)beynes,1,3-disethyl-2,3-dise

The hollow multilayer moldings of the present invention are obtained by extruding the respective resin layers described above through a multilayer die using a plurality of extruders and combine the layers before blow molding can be performed. For large vessels such as fuel tanks, drum cans and the like, drawdown resistance is more strictly required and fuel tanks are of complicated configuration since baffles are arranged therein so that local thinning tends 5 to occur and hence puncture resistance, parison thickness controllability (parison control response), pinch-off shape properties, fire resistance and the like are required. In the present invention, use of the above-mentioned polymer or composition thereof of the present invention allows these requirements to be satisfied.

Further, in the present invention, it is desirable in view of economy to use, as an exterior layer of the hollow multilayer vessels, defective moldings of a laminate containing a layer of the above-mentioned (co)polymer of the present invention or composition thereof, an adhesive layer and a barrier leyer, a recycled material such as burr, or a mixture of the recycled material, the (co)polymer of the present invention and general polyoletin resin. The blending

amounts of the recycled material and polyolefin are not limited particularly. The layer structure of the hollow multilayer vessels includes high density polyethylene (HDPE) layer of the invention/adhesive layer/polyamide resin (PA), HDPE layer of the invention/adhesive layer/EVOH, HDPE layer of the invention/adhesive layer/polyester resin (PET), HDPE layer of the invention/adhesive layer/PA/adhesive layer/PDPE layer of the invention, HDPE layer of the invention/adhesive layer/ethylene-vinyl acetale copolymer saponification product (EVOH)/adhesive layer/HDPE layer of the invention, HDPE layer of the invention/adhesive layer/PET/adhesive layer/HDPE layer of the invention, HDPE layer of the invention + HDPE/adhesive layer/PA/adhesive layer/HDPE layer of the invention, recycled layer/adhesive layer/PA/adhesive layer/recycled layer, recycled layer/adhesive layer/PA/adhesive sive layer/HDPE of the invention, recycled layer/adhesive layer/PA/adhesive layer/HDPE layer, recycled layer + HDPE/adhesive layer/PA/adhesive layer/HDPE layer, HDPE layer/adhesive layer/EVOH layer/adhesive layer/recycled

[0058] The thickness of respective layers of the hollow moldings is not limited perticularly. In the case of multilayer moldings, generally the thickness of barrier layer is selected in the range of 0.001 to 1.0 mm, and preferably 0.01 to 0.5 mm, and the thickness of adhesive layer is selected in the range of 0.001 to 1.0 mm, and preferably 0.01 to 0.5 mm. Further, the thickness of the layer of the (co)polymer of the present invention or composition thereof and the thickness of other polyoletin layer are selected in the range of 0.5 to 10 mm, and preferably 1.0 to 7 mm.

BEST MODE FOR CARRYING OUT THE INVENTION

Hereafter, the present invention will be described in more detail by referring to examples and comparative examples. However, the present invention is not be limited to the following examples.

The physical properties of polyethylene and polyethylene resin compositions of Examples 1 to 3 and Comparative Examples 1 to 4, respectively, were measured by the following test method.

[Test Method]

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According to JIS K6760. (1) Density:

According to ASTM D790. 40 (2) Flexural modulus:

According to JIS K6760 (3) HLMFR:

(4) Measurement of molecular weight distribution:

[0061] Measurement was performed using 150 Type GPC manufactured by Waters and two columns, Shodex HT-806M, and 1,2,4-trichlorobenzene (TBC) containing 0.05% by weight of 2,6-di-t-butyl-4-methylphend (BHT) as a solvent under the conditions of a sample amount of 0.8 mg/ml, a temperature of 140 °C, and a flow rate of 1 ml/min.

50 (5) Measurement of Die Swell:

"Capillograph 1C" manufactured by Toyo Seiko was used.

A sample molten at 230 °C was extruded through an ordice having a diameter of 1 mm, a length of 40 mm and an inlet angle of 90° in a region of shear rate (γ) of 5.08 to 24.32 sec⁻¹ and the diameter of the strand at the time 55 when the length of strand reached 75 mm was measured using a laser diameter measuring apparatus attached (distance between the outlet of capillary and die swell detector: 10 mm). Die swell ralio (DS) was calculated as De/Do where Ds is the diameter of a strand and Do is the diameter of an orifice. The slope a and intercept b were obtained by the method of least squares by the equation of relationship between DS and Ln(v):

 $DS = a \times Ln(\gamma)+b$.

(6) Measurement of Stress Ratio

Preparation of Sample

[0064] "Capillograph 1C" manufactured by Toyo Seiko was used.
[0065] A sample molten at 210 °C was extruded through an orifice having a diameter of 3 mm, a length of 15 mm and an inlet angle of 90° at a constant piston speed of 10 mm/min to obtain a sample.

Measurement of Stress Ratio

[0066] "Melten Rheometer" manufactured by Toyo Seiki was used. After annealing at 170 °C for 1,000 seconds, a sample was clamped by a clamp and stretch viscosity measurement was practiced at a strain speed of 0.1 sec. 1. The stress at a strain of 1,0 was defined 'of 0.0' and the maximum stress value generated until the sample was broken was defined 'ormax' and stress ratio "ormax/of 0.0" was calculated.

Measurement of a Value and Time for Breakage

20 (1067) A sample obtained in the same manner as in the case of measurement of stress ratio were measured using Meltan Rheometer' manufactured by Toyo Seiki. That is, a sample was molten at 170 °C and then clamped by a clamp and its stretch viscosity was measured under a constant tension such that initial stress (= tension/initial cross section

of sample) was 3.0x10° Pa. [0.68] Assuming that when I_0 is an initial sample diameter and I_1 is a sample diameter after t seconds, then [0.68] Assuming that when I_0 is an initial sample diameter and I_0 is a sample was broken was measured as a time for breakage.

(7) Extrusion Properties:

50 [0069] Multilayer blow molding machine "NB30" manufactured by Nippon Seikosho (main material extruder: 90 mm) was used and the extruded amount was measured at 60 rpm.

(8) Drawdown Resistance:

[0070] Multilayer blow molding machine "NB30" manufactured by Nippon Seikosho was used. A 1,600 cm-long parison weighing 8.5 kg was extruded at a resin temperature of 230 °C and after completion of the extrusion, changes in length of parison with lapse of time were measured. The parison after being shortened due to shrink back was saging as a result of drawdown by self weight and elongated. The length of parison at the time when it was extruded is defined L₀ and the time counted from the that time until the parison once shortened by shrinking took the length of L₀ again was defined retention time, Tp, and evaluation of drawdown resistance was performed.

(9) Pinch-off Shape Properties

10071] Using the above-mentioned blow molding machine and mold, 40-liter multilayer tanks (each weighing 6 kg).

45 having a layer construction of resin layer of example or comparative exampleadhesive layer (maletic arrhydride-modified high density polyethylene)/nylon-Bladhesive layer/resin layer of example or comparative example were molded. The pinch-off protions of the obtained moldings were cut vertically with respect to pinch-off weided line. Pinch-off properties were evaluated from the thickness distribution in the cross sectional plane as shown in Fig. 1(a). The thickness of the were evaluated from the thickness of the molden portion was defined "7 and the thickness of thicker portions on the both sides was defined "7, and pinch-off properties were evaluated by a ratio of t to T, VT, (Fig. 1(b) and (c)).

(10) Puncture Resistance:

[0072] Using a multilayer blow molding machine "NB30" manufactured by Nippon Seikosho (screw diameter: 90 mm, die diameter: 120 mm) and molds having the same inlet size of W1+200, W2+300) and different depths D (100, 150, 200, and 250 mm), box-shaped hollow moldings were molded and deep draw ratiu (D/W1) at which puncture of parison occurred was evaluated to thereby evaluate molding limit (puncture resistance).

(11) Tank Drop Test:

Using the above-mentioned multilayer tanks, drop tests were practiced. The drop tests were conducted by filling the tank with an aqueous 50% ethylene glycol solution to 70% of total volume and vertical drop tests were carried out at -40 °C at a height of 6 m.

(12) Parison Control Response

Using the above-mentioned blow molding machine and mold, box-shaped hollow moldings were obtained by setting a parison controller such that the thickness of a part of the middle portion of the molding was thickened at the time of molding. The parison controller used was a 25-point setting parison controller manufactured by MOOG, attached to the above-mentioned molding machine. By measuring the thickness distribution of the resultant hollow moldings, the controllability of parison controller for the thickness of a parison was measured. The thickness of the middle portion where the thickness was thickened was defined Tp and the thickness at standard point where no thickening 15 occurred was defined Ts and a ratio Tp/Ts was calculated, which was used as an index of parison control response.

(13) Bottle Fire Resistance Tests:

Using the above-mentioned blow molding machine, 300 ml bottles having a thickness of 2 mm or 3 mm were molded. Water was poured into the resultant moldings up to 50% of the inner volume and the moldings were sealed. Then the moldings were burned with propane gas and the time until a hole opened was evaluated as fire resistant time.

(14) Press Plate Melting Test:

Press plates having a thickness of 2 mm or 3 mm were molded. The resultant press plates were burned from one side thereof with a mixed gas of methane and hydrogen and the time until the other side was molten was evaluated as melting time.

Example 1

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[0077] Component 1 polymerized with a Phillips catalyst and Components 2 and 3 polymerized with MgCl₂ carried Ziegler catalyst were blended in a biaxial extruder (KTX-90 manufactured by Kobe Seikosho) to prepare a sample of Example 1. The detailed data and blend ratio for each component are shown in Table 1. The measured properties data are shown in Table 6.

Table 1

Component	Blending proportion wt%	HL & MFR g/10min	Density g/cm ³
Component 1	70	HL=1.9	0.956
		HL=0.30	0.942
Component 2	20	MFR=800	0.970
Component 3	20	HL=2.3	0.955
Example 1		HL=2.0	

Example 2

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Component 1 polymerized with a Phillips catalyst and Components 2 and 3 polymerized with MgCl₂ carried 50 Ziegler catalyst were blended in a biaxial extruder (KTX-90 manufactured by Kobe Seikosho) to prepare a sample of Example 1. The detailed data and blend ratio for each component are shown in Table 2. The measured properties data are shown in Table 6.

Table 2

			Describe alcm3
Component	Blending proportion wt%	HL & MFR g/10min	Density gran
Component 1		HL=1.9	0.956
Component .			

Table 2 (continued)

Component	Blending proportion wt%	HL & MFR g/10min	Density g/cm ³
Component 2	10	HL=0.30	0.942
Component 3		MFR=800	0.970
Example 2		HL=3.1	0.957

Example 3

[0079] Component 1 polymerized with a Phillips catalyst and Components 2 and 3 polymerized with $MgCl_2$ carried Ziegler catalyst were blended in a biaxial extruder (KTX-90 manufactured by Kobe Seikosho) to prepare a sample of Example 3. The detailed data and blend ratio for each component are shown in Table 3. The measured properties data are shown in Table 6.

Table 3

			Ittoro	
g/cm ³	Density g/cr	HL & MFR g/10min	Blending proportion wt%	Component
56	0.956	HL=1.9	65	
42	0.942	HL=0.30	10	
72	0.972	MFR=810		
58	0.958	HL=4.5		
72	0.972	HL=0.30 MFR=810	10	Component 2 Component 3 Example 3

Comparative Example 1

[0800] Component 1 polymerized with a Phillips catalyst and Components 2 and 3 polymerized with MgCl₂ carried 30 Ziegler catalyst were blended in a biaxial extruser (KTX-90 manufactured by Kobe Selioscho) to prepare a sample of Comparative Example 1. The detailed data and blend ratio for each component are shown in Table 4. The measured properties data are shown in Table 6.

Table 4

Component	Blending proportion wt%	HL & MFR g/10min	Density g/cm3
Component 1	70	HL=1.8	0.953
Component 2	10	HL=0.30	0.942
Component 3	20	MFR=790	0.968
		HL=3.0	0.954
Comparative Example 1	1		

45 Comparative Example 2

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[0081] Component 1 polymerized with a Phillips catalyst and Components 2 polymerized with MgOl₂ carried Ziegler catalyst were blended in a biaxial extruder (KTX-90 manufactured by Kobe Seikosho) to prepare a sample of Comparative Example 2. The detailed deta and blend ratio for each component are shown in Table 5. The measured properties data are shown in Table 6.

Table 5

Component	Blending proportion wt%	HL & MFR g/10min	Density g/cm3
Component 1	90	HL=1.9	0.956
Component 2	10	MFR=800	0.970

Table 5 (continued)

180le 5 (continued)									
Component	Blending proportion wt%	HL & MFR g/10min	Density g/cm ³						
Comparative Example 2		HL=4.2	0.957						
Comparative									

Comparative Example 3

[0082] Using a commercially available high density polyethylene for blow molding (density = 0.947g/cm³, HLMFR =
 4.6, JLex HD 4551H, manufactured by Japan Polyoletins Co., Ltd.) extrusion properties and the like were evaluated.
 The results are shown in Table 6.

Comparative Example 4

[0083] Using a commercially available high density polyethylene for blow molding (density = 0.951 g/cm³, HLMFR = 15 5.6, Hizex 8200B, manufactured by Mitsui Petrochemical Co., Ltd.) extrusion properties and the like were evaluated. The results are shown in Table 6.

	Comparative Example 4	0.951	13000	9 4	65 6	0.029	12	1.78	OORE	964	2	8 3	8 6	10	Direct const	1 32	8	184	24	44
	Comparative Example 3	0.947	11000	4.6	ē	0.036	1.29	3,5	0.048	16.1	10	5 6	3	>1.25		4	88	167	2	64
	Comparative Example 2	0.957	15200	5.4	8	0.094	1.40	2.97	0.033	26	æ	15	0,1	>1.25	c	1.32	109	190	31	2
	Comparative Example 1	0.954	14000	3.0	31	0.032	1.31	3.05	0.025	30.6	109	98	1.0	>1.25	0	1.50	105	185	27	48
Table 6	Example 3	0.958	16000	4.5	ĝ	0.033	1.29	2.98	0.026	30.2	120	\$	0.7	>1.25	0	1.48	18	208	38	99
	Example 2	0.957	15500	3.1	34	0.030	1.30	3.01	0.024	31.5	107	š	5.	>1.25	0	1.51	117	504	35	55
	Example 1	0.955	-	2.3	40	0.029	1.32	2.85	1	1	101	99^	5	>1.25	0	1.50		'	'	-
Ī			é.		c _o	2	æ	6		nd)) Liu	(puope	(£)		Œ	Tp/Ts)	2mm	3mm	2mm	3mm
	ITEMS.	Density (g/cm²)	Flexural modulus (Kgf/cm²)	HLMFR (g/10min)	Molecular weight distribution (Mw/Mn)	(s) edolS	Intercept (b)	Stress ratio ($\sigma \max \sigma$ 1.0)	α Vajue	Time for breakage (second)	Extrusion properties (kg/hr)	Drawdown resistance (Tp: second)	Pinch-off shape properties (t/T)	Puncture resistance (Blow ratio:D/W1)	Tank drop test (-40 °C, 6m)	Parison control response (Tp/Ts)	Bottle fire resistance test	(hole opening time: second)	Press plate melting test	(mening time: second)
	-	Den	Flexural m	HLM	Molecular	Die swell	рирешев	Stress ra		Time for b	Extrusion	Drawdown re	Pinch-off st	Punci (Blo	Tank dro	Parison con	Bottle fire re	(hole opening	Press plate	a Gumem)

INDUSTRIAL APPLICABILITY

[0084] As described above, by use of the (co)polymer of the present invention there can be obtained hollow moldings which are excellent in drawdown resistance, puncture resistance, parison control response, pinch-off shape and fire resistance, causes no local trinning and is excellent in mechanical strength such as impact strength and rigidity. It can be used advantageously as a material for blow molding of large hollow moldings such as fuel tanks and drum cans having complicated shapes such as ballles.

Claims

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- 1. An ethylene (co)polymer which satisfies the following requirements (a) to (d):
 - (a) a density of 0.93 to 0.98 g/cm³,
 - (b) a molecular weight distribution (Mw/Mn) of 25 to 50,
 - (c) that values of slope a and of intercept b obtained from a die swell (DS) and a shear rate (r) measured in a region of shear rate (r) (6.06 to 24.8 sec⁻¹) at 230 °C, in accordance with the equation (1) below

$$DS = a \times Ln(\gamma) + b \tag{1}$$

- are such that a≤0.05 and b≤1.45.
 - (d) that the stress ratio of a stress (o) at a strain of 1.0 to a maximum stress (omax) in a stress-strain curve in unlaxial stretching at 0.1 sec¹ measured at 170 °C is such that omax/o≥2.1.
- The ethylene (co)polymer as claimed in claim 1, wherein the ethylene (co)polymer further satisfies the requirement
 (e) that a value of (c) as defined by the equation (2) below

$$\alpha = \epsilon / t$$
 (2)

- (wherein α is LM when ε defined by ε = -Ln(sample diameter after t seconds I,/initial sample diameter I₀) is 0.6) when uniaxially stretched under a constant tension such that initial stress (= tension/initial cross section of sample) measured at 170 °C is 3.0x10⁴ Pa is not greater than 0.03 and time for sample breakage is not shorter than 25 seconds.
- The ethylene (co)polymer as claimed in claim 1 or 2, wherein the ethylene (co)polymer further satisfies the requirement (f) HLMFR s 10 g/10 min.
 - 4. The ethylene (co)polymer as claimed in any one of claims 1 to 3, wherein the ethylene (co)polymer has a C_{3-20} α -olefin content of 10 mol% or less.
- 5. A hollow molding comprising the ethylene (co)polymer as claimed in any one of claims 1 to 4, or composition thereof.
 - 6. The hollow molding as claimed in claim 4, wherein the hollow molding is a fuel tank.
- 45 7. A laminate comprising a layer comprising at least the ethylene (co)polymer as claimed in any one of claims 1 to 4 or compositions thereof and a barrier layer.
 - The laminate as claimed in claim 7, wherein the laminate comprises at least a layer of the ethylene (co)polymer as daimed in any one of claims 1 to 4 or compositions thereof, an adhesive layer, and a barrier layer.
 - The laminate as claimed in claim 7 or 8, wherein the barrier layer comprises at least one member selected from
 polyamide resins, ethylene/vinyl acetate copolymer saponification products, poly(meth)acrylonitrile resins, polyacetal resins, polyester resins, polyvinylidene chloride resins or compositions thereof.
- 59 10. The laminate as claimed in claim 8 or 9, wherein the adhesive layer comprises at least one adhesive resin selected from the group consisting of copolymers of unsaturated carboxylic acids or derivatives thereof with olefins and modified polyclefin resins grafted with unsaturated carboxylic acids or derivatives and mixtures of the adhesive resin with unmodified polyclefin.

- 11. A blow molded multilayer product comprising a laminate which comprises a layer comprising the ethylene (co)polymer as claimed in any one of claims 7 to 10 or compositions thereof, a barrier layer, and optionally an adhesive layer.
- 5 12. The blow molded multilayer product as claimed in claim 11, wherein the blow molded multilayer product is a fuel tank.
 - 13. A blow molded multilayer product comprising a laminate which comprises a layer of at least one resin selected from recycled resins composed of the laminate as claimed in any one of claims 7 to 10, compositions of the recycled resins and the ethylene (co)polymer as claimed in any one of claims 1 to 3, or compositions of them with polyolefin resins, a barrier layer, and optionally an achesive layer.
 - 14. The blow molded multilayer product as claimed in claim 13, wherein the blow molded multilayer product is a fuel tank.

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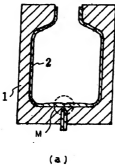
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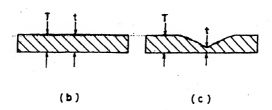
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Fig. 1





INTERNATIONAL SEARCH REPORT

ternational application No. PCT/JP98/02784

CLASSIFICATION OF SUBJECT MATTER

1 1 3 4

Int.C1 C08L23/04, C08F10/02, B32B27/32

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED nimum documentation searched (classification system followed by classification symbols)
Int. C1 C08L23/00-23/36, C08F10/00-10/14, C08F110/00-110/14,

COSF210/00-210/18

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	JP, 59-196146, A (Asahi Chemical Industry Co., Ltd.), 'November, 1984 (07. 11. 84), Claims; page 3, upper left column, line 3 to page 6, upper left column, line 15 (Family: none)	1-14
A	JP, 06-248124, A (Nippon Oil Co., Ltd.), 6 September, 1994 (06. 09. 94), Claims; Par. Nos. [0004] to [0020] (Family: none)	1-14
EA	JP, 09-169087, A (Nippon Unicer K.K.), 30 June, 1997 (30. 06. 97), Claims; Par. Nos. [0005] to [0021] (Pamily: none)	1-14
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Further documents are listed in the continuation of Box C. See patent family annex.

Special categories of cited documents: document defining the general state of the un which is not considered to be opticitable relevances cartier document but published on or after the international filling data document which any throw doubts on priority claim(4) or which is cited to entablish the publication can be of another citation or other document published prior to the international filing date but later than the priority date claimed

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council or particular resovance; we distinct invention clear asidered to involve as investive step when the document in misland with one or more other such documents, such conf-ing obvious to a person skilled in the art council member of the terms patent family

Date of mailing of the international search report 8 September, 1998 (08. 09. 98)

Name and mailing address of the ISA/ Japanese Patent Office Telephone No.

Form PCT/ISA/210 (second sheet) (July 1992)

Date of the actual completion of the international search 24 August, 1998 (24. 08. 98)

Authorized officer